

APPENDIX A
“CLEAN” VERSION OF EACH PARAGRAPH/SECTION/CLAIM
37 C.F.R. § 1.121(b)(ii) AND (c)(i)

CLAIMS (with indication of amended or new):

(Amended) 3. The pharmaceutical composition according to claim 1, wherein the stabilizing agent is saccharose alone.

(Amended) 4. The pharmaceutical composition according to claim 1, containing 3 or 10 mg/vial of hGRF.

(Amended) 5. The pharmaceutical composition according to claim 1 comprising 3 or 10 mg/vial of hGRF and 20.52 or 68.4 mg/vial of saccharose.

(Amended) 6. The pharmaceutical composition according to claim 1 further comprising buffering agents.

(Amended) 7. A process for preparing a pharmaceutical composition according to claim 1, comprising the preparation of an aqueous solution of the components, the distribution within containers and the lyophilization in the containers.

(Amended) 8. Forms of presentation of said pharmaceutical composition comprising the solid mixture according to claim 1, hermetically closed in a sterile condition within a container suited for storage before use and for reconstitution of the mixture into a solvent or into a solution for injectables.

(Amended) 9. A solution comprising the solid mixture according to claim 1, reconstituted in a solvent or a solution for injectables.